118TH CONGRESS 2D SESSION **S**.

To amend the Federal Food, Drug, and Cosmetic Act to ensure the safety of food and limit the presence of contaminants in infant and toddler food, and for other purposes.

IN THE SENATE OF THE UNITED STATES

Ms. KLOBUCHAR (for herself and Ms. DUCKWORTH) introduced the following bill; which was read twice and referred to the Committee on

A BILL

- To amend the Federal Food, Drug, and Cosmetic Act to ensure the safety of food and limit the presence of contaminants in infant and toddler food, and for other purposes.
 - 1 Be it enacted by the Senate and House of Representa-
 - 2 tives of the United States of America in Congress assembled,
 - **3** SECTION 1. SHORT TITLE.
 - 4 This Act may be cited as the "Baby Food Safety Act5 of 2024".

1 SEC. 2. DEFINITION OF INFANT AND TODDLER FOOD. 2 Section 201 of the Federal Food, Drug, and Cosmetic 3 Act (21 U.S.C. 321) is amended by adding at the end the following: 4 5 "(tt) The term 'infant and toddler food' means food that purports to be, or is represented as being, specifically 6 7 for infants or children up to the age of 24 months.". 8 SEC. 3. CONTAMINANTS IN FOOD, INCLUDING INFANT AND 9 **TODDLER FOOD.** 10 (a) IN GENERAL.—Chapter IV of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 341 et seq.) is amend-11 12 ed by adding at the end the following: 13 "SEC. 425. CONTAMINANTS IN FOOD, INCLUDING INFANT 14 AND TODDLER FOOD. 15 "(a) Administrative Orders for Contaminants IN FOOD.— 16 17 "(1) IN GENERAL.—Within the applicable time-18 frame specified in paragraph (4), the Secretary, by 19 administrative order— 20 "(A) shall establish limits on— 21 "(i) lead, cadmium, mercury, and 22 total arsenic in infant and toddler food; 23 "(ii) lead, cadmium, mercury, and 24 total arsenic in food pouches made with 25 fruit or vegetable puree or juice; and 26 "(iii) lead and arsenic in juice; and

1	"(B) if the Secretary determines appro-
2	priate upon review of relevant health data and
3	other relevant available information, may—
4	"(i) establish limits for additional con-
5	taminants in infant and toddler food;
6	"(ii) establish limits for additional
7	contaminants in juice;
8	"(iii) establish limits for additional
9	contaminants in food pouches made with
10	fruit or vegetable puree or juice; and
11	"(iv) revise limits established pursu-
12	ant to subparagraph (A).
13	"(2) PROCEDURE.—In establishing or revising
14	any limit under paragraph (1), the Secretary shall—
15	"(A) evaluate relevant health data and
16	other information the Secretary considers rel-
17	evant;
18	"(B) take into account relevant differences
19	among food types, groups, and categories, as
20	appropriate, including the extent to which the
21	presence of a contaminant cannot be avoided;
22	and
23	"(C) notwithstanding the requirements of
24	subchapter II of chapter 5 of title 5, United

1	States Code, and chapter 6 of title 5, United
2	States Code—
3	"(i) publish any administrative order
4	under paragraph (1) in the Federal Reg-
5	ister following—
6	"(I) publication of a proposed
7	order in the Federal Register; and
8	"(II) consideration of comments
9	to a public docket open for not fewer
10	than 45 calendar days; and
11	"(ii) set forth in any proposed or final
12	administrative order under paragraph (1)
13	a substantive summary of the valid sci-
14	entific evidence concerning the proposed or
15	final limit.
16	"(3) Additional contaminants; changes to
17	LIMITS.—If the Secretary determines appropriate
18	after review of relevant data and available health in-
19	formation, the Secretary may revise any limit estab-
20	lished under this subsection by administrative order
21	published in the Federal Register in accordance with
22	paragraph $(2)(C)$.
23	"(4) TIMEFRAME FOR INITIAL LIMITS.—
24	"(A) PROPOSED ORDERS.—Subject to the
25	requirements of paragraph (2)(C), the Sec-

1	retary shall issue proposed orders for limits
2	under paragraph (1)(A) as follows:
3	"(i) For lead, not later than Decem-
4	ber 31, 2025.
5	"(ii) For total arsenic, not later than
6	December 31, 2025.
7	"(iii) For cadmium, not later than
8	April 30, 2026.
9	"(iv) For mercury, not later than
10	April 30, 2028.
11	"(B) FINAL ORDERS.—The Secretary shall
12	issue each final administrative order for a limit
13	established pursuant to paragraph (1)(A) not
14	later than the earlier of—
15	"(i) the applicable deadline for a final
16	order specified in paragraph (1); or
17	"(ii) 18 months after issuance of the
18	respective proposed order.
19	"(5) CRITERIA.—The limits established under
20	this section shall represent the level at which the
21	Secretary finds necessary for the protection of public
22	health. In determining such limits the Secretary
23	shall take into account the extent to which the use
24	of such substance is required or cannot be avoided
25	in the production of each such article, and the other

ways in which a consumer may be affected by the
 same or other contaminants, taking into consider ation relevant information and data that has been
 made available.

5 "(6) ADULTERATED FOOD.—A food may be de-6 termined adulterated, at the final product stage, 7 under section 402(j), if such food bears or contains 8 any contaminant in excess of a limit established 9 under this subsection when considering variability of 10 the validated method of analysis.

11 "(7) PERIODIC REVIEW.—The Secretary shall 12 periodically review the limits established under this 13 subsection, taking into consideration relevant infor-14 mation and available data to consider whether such 15 limits should be revised, following the procedure de-16 scribed in paragraph (2), in accordance with the cri-17 teria specified in paragraph (5).

18 "(b) SAMPLING AND TESTING FOR CONTAMINANTS19 IN FOOD, INCLUDING INFANT AND TODDLER FOOD.—

"(1) IN GENERAL.—Beginning not later than
180 days after the date of enactment of the Baby
Food Safety Act of 2024, the owner, operator, or
agent in charge of a facility engaged in manufacturing or processing infant and toddler food, food
pouches made with fruit or vegetable puree or juice,

1	or juice for consumption in the United States
2	shall—
3	"(A) have a control program pursuant to
4	section 418 in place for contaminants subject to
5	ordered limits under subsection (a), or be in
6	compliance with the Juice Hazard Analysis
7	Critical Control Points Program of the Food
8	and Drug Administration, as applicable;
9	"(B) be in compliance with regulations
10	promulgated under section 420(b);
11	"(C) collect representative samples of each
12	such food in final product form in accordance
13	with a sampling plan described in paragraph
14	(2); and
15	"(D) conduct testing of the samples col-
16	lected from the final food product for contami-
17	nants, in accordance with such sampling plan.
18	"(2) Requirements for sampling plan.—
19	"(A) IN GENERAL.—The owner, operator,
20	or agent in charge of a facility described in
21	paragraph (1) shall—
22	"(i) prepare a written sampling plan
23	for all sampling and testing required under
24	this subsection; and

1	"(ii) ensure that all sampling and
2	testing conducted under this subsection is
3	conducted in accordance with the sampling
4	plan.
5	"(B) SAMPLING PLAN.—A sampling plan
6	required by subparagraph (A) shall identify—
7	"(i) the number of sampling units and
8	sample unit size based upon appropriate
9	criteria for identifying, in a representative
10	fashion, the levels of contaminants in each
11	food; and
12	"(ii) one or more appropriate test
13	methods and procedures to be used to ana-
14	lyze the samples.
15	"(C) GUIDANCE.—Not later than 18
16	months after the date of enactment of the Baby
17	Food Safety Act of 2024, the Secretary shall
18	issue guidance to assist facilities described
19	under paragraph (1) with developing sampling
20	plans. Such guidance may, as the Secretary de-
21	termines appropriate, address when samples
22	should be tested for specific species of contami-
23	nants.
24	"(3) Contaminants to be tested.—In car-
25	rying out the sampling and testing under this sub-

1	section, the owner, operator, or agent in charge of
2	a facility described in paragraph (1) shall ensure
3	that each sample is tested for levels of—
4	"(A) lead, cadmium, mercury, and total ar-
5	senic;
6	"(B) any other contaminant that the Sec-
7	retary may specify by regulation, and in accord-
8	ance with the sampling plan under paragraph
9	(2).
10	"(4) Foods to be tested.—The sampling
11	and testing conducted under this subsection shall be
12	conducted for—
13	"(A) infant and toddler foods, in final
14	product form;
15	"(B) pouches made with fruit and vege-
16	table puree or juice;
17	"(C) juice; and
18	"(D) such other foods in final product
19	form as the Secretary may specify, by regula-
20	tion, as appropriate to protect the public health.
21	"(5) Recordkeeping.—
22	"(A) IN GENERAL.—The owner, operator,
23	or agent in charge of a facility described in
24	paragraph (1) shall maintain, for not less than
25	2 years or the shelf-life of each food product

1	manufactured or processed by the facility,
2	whichever is longer, records documenting the
3	sampling plan and results of testing conducted
4	under this subsection with respect to the food.
5	The owner, operator, or agent in charge of such
6	a facility shall make such records available for
7	inspection by the Secretary upon request by the
8	Secretary.
9	"(B) REQUIREMENTS.—The records main-
10	tained as required under subparagraph (A)
11	shall include—
12	"(i) a detailed description of the foods
13	sampled and tested;
14	"(ii) the number of samples and tests
15	performed;
16	"(iii) the size and number of items in
17	each sample unit;
18	"(iv) a copy of the sampling plan re-
19	quired under paragraph (2);
20	"(v) identification of the entity con-
21	ducting the sampling;
22	"(vi) identification of the entity con-
23	ducting the testing; and

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1	"(vii) identification of the analytical
2	methods used to perform the sampling and
3	testing.
4	"(C) Applicability.—The requirements
5	of this paragraph shall apply to all records of
6	sampling and testing conducted pursuant to
7	this subsection, regardless of the findings.
8	"(6) LABORATORY ACCREDITATION.—The
9	owner, operator, or agent in charge of a facility de-
10	scribed in paragraph (1) shall ensure that testing re-
11	quired pursuant to this subsection is performed in
12	accordance with international standards by a labora-
13	tory that is accredited by an accreditation body that
14	conforms to international accreditation standards.
15	Testing conducted under this subsection is not sub-
16	ject to the requirements regarding laboratory accred-
17	itation described in section 422.
18	"(7) Sampling and testing program.—The
19	Secretary shall develop and implement a sampling
20	and testing program for infant and toddler food for
21	sale to consumers that is sufficient to—
22	"(A) support the periodic review under
23	subsection (a)(7) of limits on lead, cadmium,
24	mercury, and arsenic in infant and toddler food;
25	and

1	"(B) independently verify the effectiveness
2	of the sampling and testing conducted pursuant
3	to this subsection by the owner, operator, or
4	agent in charge of a food facility.
5	"(8) GUIDANCE.—The Secretary shall issue
6	guidance to assist food facilities in complying with
7	this subsection.
8	"(c) Record Availability.—
9	"(1) IN GENERAL.—Upon request by the Sec-
10	retary, the owner, operator, or agent in charge of a
11	facility described in subsection (b)(1) shall—
12	"(A) make all records required under this
13	section available promptly to the Secretary for
14	inspection and copying; and
15	"(B) provide within a reasonable time an
16	English translation of such records maintained
17	in a language other than English.
18	"(2) Record availability in Lieu of an in-
19	SPECTION.—Any records that the Secretary may in-
20	spect under this section shall, upon the request of
21	the Secretary, be provided to the Secretary by the
22	owner, operator, or agent in charge of a facility de-
23	scribed in subsection $(b)(1)$, in advance of or in lieu
24	of an inspection, within a reasonable timeframe,
25	within reasonable limits, and in a reasonable man-

ner, and in either electronic or physical form, at the
 expense of such owner, operator, or agent. The Sec retary's request shall include a sufficient description
 of the records requested.

5 "(3) CONFIRMATION.—Upon receipt of records 6 requested under paragraph (1) or (2), the Secretary 7 shall provide to the owner, operator, or agent de-8 scribed in paragraph (2) confirmation of the receipt. 9 "(4) AUTHORITY OF THE SECRETARY.-Noth-10 ing in this subsection supplants the authority of the 11 Secretary to conduct sampling, testing, or inspec-12 tions otherwise permitted under this Act in order to ensure compliance with this Act. 13

14 "(d) DELAYED APPLICABILITY.—The requirements
15 for sampling and testing under this section shall apply be16 ginning on the date that is 2 years after the date of enact17 ment of this subsection.

18 "(e) PREEMPTION OF STATE AND LOCAL REQUIRE-19 MENTS REGARDING FOOD INGREDIENTS AND CONTAMI-20 NANTS IN FOOD, INCLUDING INFANT AND TODDLER 21 FOOD.—No State or political subdivision of a State may 22 establish or continue in effect with respect to contami-23 nants in food, including infant and toddler food, food 24 pouches made with fruit or vegetable pure or juice, and 25 juice, any requirement that is different from, or in addi-

tion to, or not identical with any requirement under this
 section, and relates to contaminant sampling and testing,
 contaminant limits, disclosure of contaminant test results,
 contaminant labeling, contaminant warnings, or any other
 matter related to contaminants in food.".

6 (b) IMPORTER REQUIREMENTS.—Section 805(c)(4)
7 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C.
8 384a(c)(4)) is amended, by inserting ", including as de9 scribed in section 425(b)" before the period at the end.
10 (c) ENFORCEMENT.—

(1) ADULTERATION.—Section 402 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 342)
is amended by adding at the end the following:

14 "(j) If it is an article of food in final product form
15 that is an infant and toddler food, a food pouch made with
16 fruit or vegetable puree or juice, or juice and—

17 "(1) such food bears or contains any contami18 nant in excess of limits established under section
19 425(a); or

"(2) the owner, operator, or agent in charge of
a facility that manufactures or processes the food is
not in compliance with subsection (b) or (c) of section 425.".

(2) PROHIBITED ACT.—Section 301 of the Fed eral Food, Drug, and Cosmetic Act (21 U.S.C. 331)
 is amended by adding at the end the following:

4 "(jjj) The failure of an owner, operator, or agent in
5 charge of a facility that manufactures or processes food
6 to comply with applicable requirements under subsection
7 (b) or (c) of section 425.".

8 SEC. 4. IMPLEMENTATION OF FOOD TRACEABILITY PLAN; 9 STUDY ON INSPECTIONS; REPORTING ON IN10 SPECTIONS.

11 IMPLEMENTATION PLAN.—The Secretary of (a) 12 Health and Human Services (referred to in this section 13 as the "Secretary"), acting through the Commissioner of 14 Food and Drugs, in coordination with the FDA Human 15 Foods Program and the Center for Food Safety and Applied Nutrition, shall finalize an implementation plan for 16 17 the Food and Drug Administration to achieve its goal of compliance, not later than January 20, 2026, with the rule 18 issued by the Food and Drug Administration titled, "Re-19 20 quirements for Additional Traceability Records for Cer-21 tain Foods" (87 Fed. Reg. 70910 (November 21, 2022)). 22 Such plan shall include a description of—

23 (1) any resource needs of the Food and Drug24 Administration;

1	(2) strategies for facilitating compliance with
2	the rule; and
3	(3) detailed plans for communicating with and
4	educating regulated entities, non-Federal regulatory
5	partners, and regulatory staff of the Food and Drug
6	Administration about the requirements under the
7	rule.
8	(b) STUDY ON INSPECTIONS.—The Secretary shall—
9	(1) conduct a study to—
10	(A) determine the annual number of facil-
11	ity inspections that is sufficient to determine
12	that imported foods are held to the same safety
13	standards as domestic food; and
14	(B) identify whether such inspection tar-
15	gets are consistent with the targets in the most
16	recent annual report regarding food conducted
17	under section 1003(h) of the Federal Food,
18	Drug, and Cosmetic Act (21 U.S.C. 393(h));
19	and
20	(2) not later than 1 year after the date of en-
21	actment of this Act, submit a report to Congress on
22	the findings of such study, and, if applicable, any
23	factors preventing the Secretary from meeting its
24	goal for the number of inspections and a plan to en-
25	sure that such goal is met in the next 2 years.

1	(c) ANNUAL REPORT REGARDING FOOD.—Section
2	1003(h)(1) of the Federal Food, Drug, and Cosmetic Act
3	(21 U.S.C. 393(h)(1)) is amended—
4	(1) in subparagraph (E), by striking "and" at
5	the end;
6	(2) in subparagraph (F), by striking the period
7	and inserting "; and"; and
8	(3) by adding at the end the following:
9	"(G) the nature of domestic facility and
10	foreign facility inspections described in subpara-
11	graph (C), the aggregate inspection findings of
12	such inspections, and the compliance rate of
13	foreign food importers with certification stand-
14	ards;".
14	
	SEC. 5. RECORDS FOR OR IN LIEU OF CERTAIN INSPEC-
14 15 16	SEC. 5. RECORDS FOR OR IN LIEU OF CERTAIN INSPEC- TIONS.
15 16	
15	TIONS.
15 16 17	TIONS. Section 704(a)(4) of the Federal Food, Drug, and
15 16 17 18	TIONS. Section 704(a)(4) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 374(a)(4)) is amended—
15 16 17 18 19	TIONS. Section 704(a)(4) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 374(a)(4)) is amended— (1) by redesignating subparagraphs (B)
15 16 17 18 19 20	TIONS.Section 704(a)(4) of the Federal Food, Drug, andCosmetic Act (21 U.S.C. 374(a)(4)) is amended—(1) by redesignating subparagraphs (B)through (D) as subparagraphs (C) through (E), re-
 15 16 17 18 19 20 21 	TIONS. Section 704(a)(4) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 374(a)(4)) is amended— (1) by redesignating subparagraphs (B) through (D) as subparagraphs (C) through (E), re- spectively;
 15 16 17 18 19 20 21 22 	TIONS. Section 704(a)(4) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 374(a)(4)) is amended— (1) by redesignating subparagraphs (B) through (D) as subparagraphs (C) through (E), re- spectively; (2) by inserting after subparagraph (A) the fol-

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1 a person that owns or operates, or is an agent in charge 2 of, an establishment that is engaged in any of the activi-3 ties described in clause (ii) shall, upon the request of the 4 Secretary, be provided to the Secretary by such person, 5 in advance of or in lieu of an inspection, within a reasonable timeframe, within reasonable limits, and in a reason-6 7 able manner, and in either electronic or physical form, at 8 the expense of such person. The Secretary's request shall 9 include a sufficient description of the records requested. 10 "(ii) The activities described in this clause are 11 records relating to—

"(I) the manufacturing, processing, packing,
transporting, distributing, receiving, holding, or importing of an article of food; or

"(II) the distribution or use of animal feed
bearing or containing a veterinary feed directive
drug, or the issuance of a veterinary feed directive.";
and

19 (3) by adding at the end the following:

20 "(F) Section 703 does not apply to records or other
21 information obtained pursuant to a request made under
22 this section.".

23 SEC. 6. MANDATORY RECALL AUTHORITY.

Section 423(a) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 350l(a)) is amended by inserting

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after "animals," the following: "or if the Secretary deter mines through any means that an article of infant and
 toddler food (other than infant formula) bears or contains
 a contaminant that renders the product adulterated under
 section 402(a)(1),".

6 SEC. 7. ENVIRONMENTAL MONITORING.

7 Chapter IV of the Federal Food, Drug, and Cosmetic
8 Act (21 U.S.C. 341 et seq.), as amended by section 3,
9 is further amended by adding the following:

10"SEC. 426. ENVIRONMENTAL MONITORING OF INFANT AND11TODDLER FOOD.

12 "(a) IN GENERAL.—A manufacturer of infant and 13 toddler food shall establish and implement an environmental monitoring program to verify the effectiveness of 14 15 sanitation and hygiene controls where the food has the potential to be exposed to environment pathogens during the 16 17 manufacturing and packing process. The environmental monitoring program shall be written and include proce-18 19 dures for determining sample location, number of samples 20 to be taken, and timing and frequency of sample collection 21 and testing.

"(b) ORGANISMS SAMPLED.—The environmental
monitoring program under subsection (a) shall include
testing for environmental pathogens, lead, arsenic, mercury, or a reliable indicator organism.

"(c) SAMPLING LOCATION AND NUMBER OF SAM PLES.—A manufacturer of infant and toddler food shall
 ensure that the sampling locations from which samples
 will be taken, and the number of sites to be tested during
 routine environmental monitoring are adequate to deter mine whether sanitation and hygiene controls are effective.

7 "(d) TIMING AND FREQUENCY.—The timing and fre8 quency for collecting and testing samples shall be ade9 quate to determine whether sanitation and hygiene con10 trols are effective.

11 "(e) RECORDS.—

12 "(1) AVAILABILITY TO THE SECRETARY.—A
13 manufacturer shall make all the records required
14 under this section available promptly to the Sec15 retary, upon request, for inspection and copying.

"(2) MAINTENANCE.—Records of environmental
monitoring conducted pursuant to this section shall
be established and maintained by the manufacturer
for not less than 2 years or the shelf-life of the food,
whichever is longer.

21 "(3) CONDITIONS OF INSPECTION.—Any
22 records that the Secretary may inspect under this
23 section shall, upon the request of the Secretary, be
24 provided to the Secretary by the manufacturer, in
25 advance of or in lieu of an inspection, within a rea-

sonable timeframe, within reasonable limits, and in
 a reasonable manner, and in either electronic or
 physical form, at the expense of such manufacturer.
 The Secretary's request shall include a sufficient de scription of the records requested.

6 "(4) CONFIRMATION OF RECEIPT.—Upon re7 ceipt of the records requested under paragraph (3),
8 the Secretary shall provide to the manufacturer con9 firmation of receipt.

"(f) AUTHORITY OF THE SECRETARY.—Nothing in
this section supplants the authority of the Secretary to
conduct inspections otherwise permitted under this Act in
order to ensure compliance with this Act.

14 "(g) EFFECTIVE DATE.—The requirements of this
15 section shall apply beginning on the date that is 2 years
16 after the date of enactment of the Baby Food Safety Act
17 of 2024.

18 "(h) RULE OF CONSTRUCTION.—Nothing in this sec19 tion shall be construed to exempt any manufacturer from
20 the requirements of this Act, including the requirements
21 under section 418.".